## Chemistry

THE STRUCTURE OF PROTEINS: TWO HYDROGEN-BONDED

HELICAL CONFIGURATIONS OF THE POLYPEPETER CHAIN

By Linus Pauling, Robert B. Corey, and H. R. Branson\*

Cates and Crellin Laboratories of Chemistry,

California Institute of Technology, Pasadena, California

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During the pest fifteen years we have been attacking the roblem of the structure of proteins in several ways. One of these ways is the complete and occurate determination of the crystal structure of amino acids, peptides, and other simple substances related to proteins, in order that information about interatomic distances, band angles, and other configurational parameters might be obtained that would permit the reliable prediction of reasonable configurations for the polypeptide chain. We have now used this information to construct two reasonable hydrogen-bonded helical configurations for the polypeptide chain; we think that it is likely that these configurations constitute an important part of the structure of both fibrous and globular proteins, as well as of synthetic polypeptides. A letter amouncing their discovery was published last year.

The problem that we have set ourselves is that of finding all hydrogen-bonded structures for a single polypeptide chain, in which the residues are equivalent (except for the differences in the side chain R). An amino acid residue (other than glycine) has no symmetry alements. The general operation of conversion of one residue of a single chain into a second residue equivalent to the first is accordingly a retation about an axis accompanied by translation along the exist. Hence the only configurations for a chain competible with our

postulate of equivalence of the residues are halical configurations. For rotational angle 180° the helical configurations may degenerate to a simple chain with all of the principal atoms, C, C' (the methyl carbon), N, and O, in the same plane.

We assume that, because of the resonance of the double bond between the

carbon-oxygen and carbon-nitrogen positions, the configuration of each residue N-C is planar. This structural feature has been verified for each of the saides that we have studied. Moreover, the resonance theory is now so well grounded and its experimental substantiation so extensive that there can be no doubt whatever about its application to the faide group. The observed C-N distance, 1.32 a, corresponds to nearly fifty percent double-bond character, and we may conclude that rotation by as much as 10° from the planer configuration would result in instability by several keal mole. The interatomic distances and bond angles within the residue are assumed to have the values shown in Figure 1. These values have been formulated by consideration of the experimental values found in the crystal structure studies of DL-slanine, 3 L-threonine, 4 N-acetylglycine, and 3-glycylglycine that have been made in our Laboratories. It is further assumed that each nitrogen atom forms a hydrogen bond with an oxygen atom of another residue, with the nitrogen-exygen distance equal to 2.72 %, and that the vector from the nitrogen atom to the hydrogen-bonded oxygen atom lies not more than 30° from the N-H direction. The energy of an N-H...O=-C hydrogen bond is of the order of 8 kcal. nole, and such great instability would result from the failure to form those bonds that we may be confident of their presence. The N-H .. O distance cannot be expected to be exactly 2.72 A, but might deviate somewhat from this value.

Solution of this problem shows that there are five and only five configurations for the chain that satisfy the conditions other than that of direction of the hydrogen bond relative to the N-H direction. These correspond to the values 165°, 120°, 108°, 98°, and 70.8° for the for the rotational angle. In the first, third, and fifth of these structures the 00 group is negatively and the N-H group positively directed along the helical axis, taken as the direction corresponding to the sequence -CHR-CO-NH-CHR- of stons in the peptide chain, and in the other two their directions are reversed. The first three of the structures are unsatisfactory, in that the N-H group does not extend in the direction of the oxygen atom at 2.72 A; the fourth and fifth are satisfactory, the angle between the N-II vector and N-O vector being about 10° and 25° for these two structures, respectively. The fourth structure has 3.69 amino acid residues per turn in the helix, and the fifth structure has 5.09 residues per turn. In the fourth structure each saide group is hydrogen-bonded to the third failde group beyond it along the helix, and in the fifth structure each is bonded to the fifth mide group beyond it; we shall call those structures either the 3.7-residue structure and the 5.1-residue structure, respectively, or the third-fadde hydrogen-bonded structure and the fifth-daide hydrogen-londed structure.

The cylindrical coordinates of the atoms of one residue for each of the two structures are given in Table I, and Dravin a and photographs of the two structures are shown in Figures 2, 3, 4, and 5.

For glycine both the 3.7-residue helix and the 5.1-residue helix could occur with either a positive or a negative rotational translation; that is, as either a positive or a negative helix, relative to the positive direction of the helical existive by the sequence of atoms in the peptide chain. For other ardno acids with the L configuration, however, the positive helix and the negative helix would differ in the position of the side chains, and it might well be expected that in each case one sense of the helix would be more stable than the other. An arbitrary assignment of the R groups has been made in Figures, 2 and 3.

The translation along the holical axis in the 3.7-residue belix is 1.47 Å, and that in the 5.1 residue belix is 0.96 Å. The values for one complete turn are 5.44 Å and 4.88 Å, respectively. These values are calculated for the hydrogen-bond distance 2.72 Å; they would have to be increased by a few percent, in case that a larger hydrogen-bond distance (2.80 Å, say) were present.

The stability of our belief structures in a non-crystalline phase depends solely on interactions between adjacent residues, and does not require that the number of residues per turn be a ratio of small integers. The value 3.69 residues per turn, for the third-smide hydrogen-bonded belix, is most closely approximated by 48 residues in thirteen turns (3.693 residues per turn), and the value 5.69 for the other belix is most closely approximated by 56 residues in case that turns. It is to be expected that the number of residues per turn would be affected somethat by change in the hydrogen-loud distance, and also that the interaction of belief molecules with neighboring similar molecules in a crystal would cause small torques in the heliges, deforming them all btly into configurations with a rational number of residues per turn. For the third-smide hydrogen-bonded belix the simplest structures of this port that we would predict are the II-residue 3-turn helix (3.67 residues per turn), the 15-residue 4-turn

helix (3.75), and the 18-residue 5-turn helix (3.60). We have found some evidence indicating that the first and third of these slight variants of this helix exist in crystalline polypeptides.

These helical structures have not previously been described. In addition to the extended polypeptide chain configuration, which for nearly thirty years has been assumed to be present in stretched hair and other proteins with the \$keratin structure, configurations for the polypoptide chain have been proposed by astbury and Bell, and especially by Huggins and by Bragg, Kendrew, and Peruts. Huggins discussed a number of structures involving intra-colecular hydrogen bonds, and Bragg, Kendrow, and Peruts extended the discussion to include additional structures, and investigated the compatibility of the structures with x-ray diffraction data for hemoglobin and myoglobin. None of these authors proposed either our 3.7-residue helix or our 5.1-residue helix. On the other hand, we would eliminate, by our basic postulates, all of the structures proposed by them. The reason for the difference in results obtained by other investigators and by us through essentially similar arguments is that both Bragg and his collaborators and Huggins discussed in detail only helical structures with an integral number of residues per turn, and no seover as a mend only a rough approximation to the regularments about interatomic distances, bond angles, and planerity of the conjugated saide group, as given by our investigations of sinplor substances. We contend that these stereochemical features must be very clasely retained in stable configurations of polypoptide chains in proteins, and that there is no special stability associated with an integral number of realdnes per turn in the helical molecule. Brugg, Kendraw, and Perutz have described a structure topologically similar to our 3.7-residue helix as a hydrogen-bonded helix with A residues per turn. In their thorough comparison of their models with Patterson projections for hemoglobin and myoglobin they eliminated this

structure, and drew the cautious conclusion that the evidence favors the non-helical 3-residue folded a-keratin configuration of astbury and Bell, in which only one third of the carbonyl and main groups are involved in intra-molecular hydrogen-bond formation.

It is our opinion that the structure of effectin, effectin, and similar fibrous proteins is closely represented by our 3.7-residue helix, and that this helix also constitutes an important structural feature in hemoglobin, myoglobin, and other globular proteins, as well as of synthetic polypeptides. We believe that the 5.1-residue helix is represented in nature by supercontracted verstin and supercontracted myosin. The evidence leading us to these conclusions will be presented in later papers.

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Summary. - Two hydrogen-bonded helical structures for a polypeptide chain have been found in which the residues are stereochemically equivalent, the interatomic distances and bond angles have values found in amino acids, poptides, and other simple substances related to proteins, and the conjugated Acide system is planar. In one structure, with 3.7 residues per turn, each carbonyl and imino group is attached by a hydrogen bond to the complementary group in the third daide group removed from it in the polypeptide chain, and in the other structure, with 5.1 residues per turn, each is bonded to the firth faide group.

## References

\*Present address, Howard University, Washington, D.C.

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1 Pauling, L., and Core, R.B., J. an. Chem. Soc., 72, 5349 (1950).

<sup>2</sup>Corey, R.B., and Donolme, J., J. Am. Chem. Soc., 72, 2899 (1950).

<sup>3</sup>Lovy, R.a., and Gorey, R.B., <u>1bid.</u>, <u>63</u>, 2095 (1941); Donohue, J., <u>1bid.</u>, <u>72</u>, 949 (1950).

4Shockeler, D.P., Donohue, J., Schomeler, V., and Corny, R.D., <u>ibid</u>, <u>72</u>, 2328 (1950).

<sup>5</sup>Carpenter, G.B., and Donohue, J., <u>ibid.</u>, 72, 2315 (1950).

Hughes, E.W., and Moore, W.J., 1616., 71, 2618 (1949).

7 sthury, W.T., and Bell, F.O., Meture, 1/7, 696 (19/1).

8 Huggins, M.L., Chem. Rev., 32, 195 (1943).

9 Bragg, L., Kendrow, J.C., and Perutz, M.F., Proc. Roy. Soc., 2203, 321 (1950).

## Legends for Figures

- Fig. 1. Dimensions of the polypeptide chain.
- Fig. 2. The helix with 3.7 residues per turn.
- Fig. 3. The holix with 5.1 residues per turn.
- Fig. 4. Plan of the 3.7-residue helix.
- Fig. 5. Plan of the 5.1-residue helix.

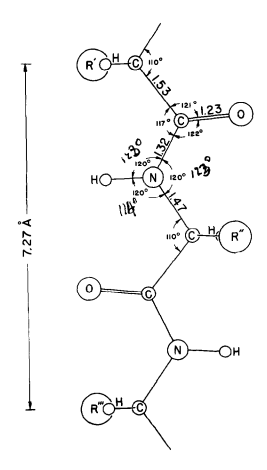


Fig. 1. Dimensions of the polypeptide chain.

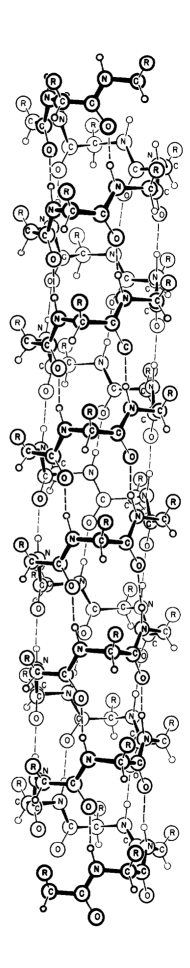


Fig. 3

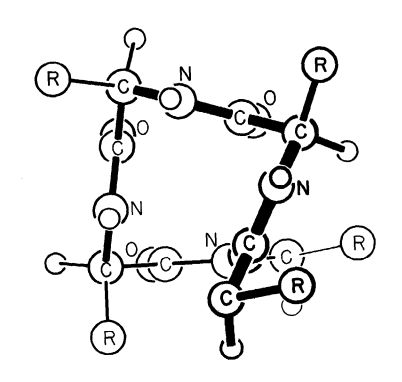


Fig. 4. Plan of the 3.7-residue helix.

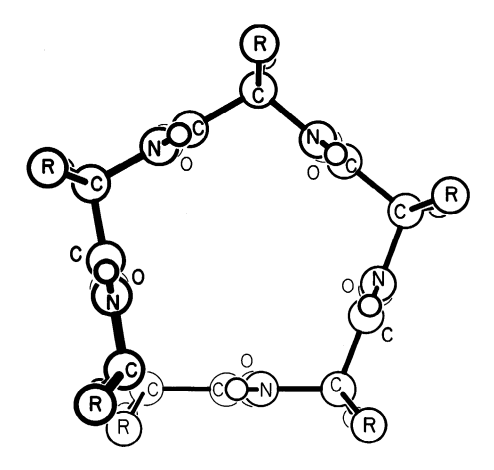


Fig. 5. Plan of the 5.1-residue helix.